Synapse–Glia Interactions at the Mammalian Neuromuscular Junction

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Perisynaptic Schwann cells (PSCs) play critical roles in regulating and stabilizing nerve terminals at the mammalian neuromuscular junction (NMJ). However, although these functions are likely regulated by the synaptic properties, the interactions of PSCs with the synaptic elements are not known. Therefore, our goal was to study the interactions between mammalian PSCs in situ and the presynaptic terminals using changes in intracellular Ca$^{2+}$ as an indicator of cell activity. Motor nerve stimulation induced an increase in intracellular Ca$^{2+}$ in PSCs, and this increase was greatly reduced when transmitter release was blocked. Furthermore, local application of acetylcholine induced Ca$^{2+}$ responses that were blocked by the muscarinic antagonist atropine and mimicked by the muscarinic agonist muscarine. The nicotinic antagonist α-bungarotoxin had no effect on Ca$^{2+}$ responses induced by acetylcholine. Local application of the cotransmitter ATP induced Ca$^{2+}$ responses that were unaffected by the P2 antagonist suramin, whereas local application of adenosine induced Ca$^{2+}$ responses that were greatly reduced by the A1 receptor antagonist 8-cyclopentyl-1,3-dimethylxanthine (CPT). However, the presence of the A1 antagonist in the perfusate did not block responses induced by ATP. Ca$^{2+}$ responses evoked by stimulation of the motor nerve were reduced in the presence of CPT, whereas atropine almost completely abolished them. Ca$^{2+}$ responses were further reduced when both antagonists were present simultaneously. Hence, PSCs at the mammalian NMJ respond to the release of neurotransmitter induced by stimulation of the motor nerve through the activation of muscarinic and adenosine A1 receptors.

Key words: adenosine; acetylcholine; Ca$^{2+}$; perisynaptic Schwann cell; transmitter release; neuromuscular junction; synapse–glia interactions

Recent evidence indicates that there are dynamic, bidirectional interactions between glial cells and neurons. Indeed, not only are glial cells modulated by nerve-evoked transmitter release but they, in turn, modulate neuronal activity (Nedergaard, 1994; Parpura et al., 1994; Pfrieger and Barres, 1997; Bezzi et al., 1998; Newman and Zaho, 1998; Robitaille, 1998).

The vertebrate neuromuscular junction (NMJ) is an interesting preparation to study synapse–glia interactions because it is a simple synapse where the anatomical relationship between presynaptic, postsynaptic, and glial elements is maintained. Perisynaptic Schwann cells (PSCs) or terminal Schwann cells, glial cells at this synapse, have been studied at the amphibian NMJ and have been shown to detect synaptic activity, as revealed by an elevation in intracellular calcium (Jahromi et al., 1992; Robitaille, 1995). PSCs possess purinergic and muscarinic receptors as well as neurokinin-1 (NK-1) receptors for substance P (Robitaille, 1995; Robitaille et al., 1997; Bourque and Robitaille, 1998), and their activity is governed by these receptors (Georgiou et al., 1994, 1999a; Bourque and Robitaille, 1998). Moreover, as a consequence of synaptic activity, PSCs modulate the efficacy of the synapse by regulating transmitter release (Robitaille, 1998; Castonguay and Robitaille, 2001).

PSCs also modulate nerve growth and synapse maintenance. Although recent evidence has been obtained at the amphibian NMJ (Herrera et al., 2000; Koirala et al., 2000), most evidence has been obtained at the mammalian NMJ. For instance, processes extended by PSCs after denervation influence the re-growth of motor axons to the muscles and their guidance back to denervated NMJs (Son et al., 1996). Furthermore, immunocytochemical markers specific for Schwann cells, synapses, and motor axons were used to show that motor axons navigate along the processes extended by PSCs from the damaged nerve endings (Son and Thompson, 1995a,b). The guidance of axons by PSCs lead to clustering of motor units, a phenomenon typical of many neurogenic pathologies (Son et al., 1996). Notwithstanding the functional importance of PSCs, very little is known about their properties at the mammalian NMJ where, in fact, no evidence of direct modulation of synaptic activity is available and where the properties are not characterized. Moreover, although synapse–glia interactions appear to be a common phenomenon at chemical synapses (Araque et al., 1999; Castonguay et al., 2001), it is still unclear how much the properties of these interactions differ between related synapses. Hence, this study aimed to characterize PSCs properties at the mammalian NMJ to determine functional similarities and differences with the properties of the amphibian NMJ.

We report that, similar to PSCs at amphibian NMJ, PSCs at the mammalian NMJ respond to the release of neurotransmitters induced by stimulation of the motor nerve via activation of muscarinic receptors. However, the role of the purinergic recep-
tors of PSCs at the mammalian NMJ differs where A1 adenosine receptors, but not P2 receptors, contribute to the activation of PSCs. Hence, these results indicate that although synapse–glia interactions are present at chemical synapses in situ, these interactions are governed by specialized features related to the synapse with which glial cells are associated.

MATERIALS AND METHODS

Nerve–muscle preparation. Experiments were performed at room temperature on levator auris longus nerve–muscle preparations dissected from CD1 mice (22–24 gm; Charles River Laboratories, Wilmington, MA). Muscles were removed under deep anesthesia (midazolam and hypnorm dissolved in water). The normal Ringer’s solution contained (in mM): 124 NaCl, 5 KCl, 1.25 NaH₂PO₄, 2 MgCl₂, 2 CaCl₂, 26 NaHCO₃, and 10 glucose. A 0 Ca²⁺/5 mM Mg²⁺ physiological solution contained (in mM): 124 NaCl, 5 KCl, 1.25 NaH₂PO₄, 5 MgCl₂, 26 NaHCO₃, and 10 glucose. Solutions were oxygenated with a gas mixture of 95% O₂ and 5% CO₂.

Calcium imaging of mammalian PSCs. Muscles were incubated for 90 min at room temperature in a physiological solution saturated with 95% O₂ and 5% CO₂. Stimulation of the motor nerve. Nerve–muscle preparations were processed for fluo-3 AM loading as described above. Stimulation of the motor nerve was achieved by direct intracellular injection of 20 μM fluo-3 AM (Molecular Probes, Eugene, OR; Tsien, 1989), 0.02% pluronic acid (Molecular Probes), and 0.5% dimethylsulfoxide (Sigma, St. Louis, MO). Muscles were then pinned down in a recording chamber coated with Silgard. Partial chelation of heavy metal ions was achieved by using 0.5 mM tetrakis(2-pyridylmethyl) ethylenediamine (Molecular Probes) to limit binding of these ions to fluo-3. Changes in fluorescence intensity were monitored using a Bio-Rad (Hercules, CA) 600 laser-scanning confocal microscope equipped with an argon ion laser. The 488 nm laser line was attenuated to 1% intensity, and a long-pass filter with a cutoff at 515 nm was used to detect the emitted fluorescence. A 40× water immersion lens was used (0.75 NA; Olympus, Tokyo, Japan). Surface NMJs were located using transmitted light microscopy. The intensity of fluorescence (F) was measured over the area of the PSCs’ cell body, and the relative changes in fluorescence intensity were expressed as % ΔF/F (F − F_rest)/F_rest × 100.

Drug applications. ATP, adenosine, muscarine, and acetylcholine (20 μM) were dissolved in the same saline as that in the recording chamber. Drugs were applied directly by micropipette (5–10 PSI, pulse duration, 200 msec) to selected cells with a micropipette (tip diameter, 2–4 μM) using a Picospritzer II (General Valve, Fairfield, NJ). The local application of the agonists induced a slight movement of the muscle fiber that moved briefly out of focus resulting in a small deflection in the fluorescence level (see Figs. 3–7). The micropipette was positioned near the PSC soma under visual control at high magnification (40× water immersion objective). There was no increase in fluorescence when Ringer’s solution was applied alone. When several applications were performed on the same cell, a recovery period of at least 15 min was allowed between each application. For the Ca²⁺–free/5 mM Mg²⁺ experiments, preparations were perfused in the modified physiological solution for 25–30 min before the experiments. Some antagonists (α-bungarotoxin, α-conotoxin MVIIIC) were applied directly in the recording chamber in a closed bath, whereas suramin, 8-cyclopentyl-1,3-dimethylxanthine (CPT), and atropine (Molecular Probes) to limit binding of these ions to fluo-3. Changes in fluorescence intensity were monitored using a Bio-Rad (Hercules, CA) 600 laser-scanning confocal microscope equipped with an argon ion laser. The 488 nm laser line was attenuated to 1% intensity, and a long-pass filter with a cutoff at 515 nm was used to detect the emitted fluorescence. A 40× water immersion lens was used (0.75 NA; Olympus, Tokyo, Japan). Surface NMJs were located using transmitted light microscopy. The intensity of fluorescence (F) was measured over the area of the PSCs’ cell body, and the relative changes in fluorescence intensity were expressed as % ΔF/F (F − F_rest)/F_rest × 100.

Stimulation of the motor nerve. Nerve–muscle preparations were processed for fluo-3 AM loading as described above. Stimulation of the distal end of the cut motor nerve (50 Hz, 30 sec) was achieved using a suction electrode. To prevent muscle contractions evoked by transmitter release, cholinergic receptors were blocked with α-bungarotoxin (α-BuTx; 20 μM; Molecular Probes). Preparations were allowed to rest for 20–25 min between trains of stimuli when several trains were performed on the same preparation.

Immunocytochemistry. After some experiments, muscles were fixed using 4% paraformaldehyde for 10 min and rinsed for at least 30 min in three changes of Ringer’s solution (20°C). Muscles were then permeabilized in −20°C methanol for 6 min, washed again in Ringer’s solution for 30 min, and nonspecific staining was blocked using Ringer’s solution containing 0.3% Triton X-100 and 0.2% dry milk. Muscles were incubated overnight at room temperature with a rabbit antibody solution raised against cow S-100 (Dako, Carpinteria, CA) prepared in the solution indicated above (dilution 1:400). After incubation in the primary antibody, muscles were rinsed in three changes of Ringer’s solution and incubated in secondary antibody for 1 hr at room temperature. The secondary antibody was an anti-rabbit conjugated to biotin-SP (Jackson ImmunoResearch, West Grove, PA: 1:500) and revealed using avidin FITC (1:100; for 2 hr). Finally, muscles were rinsed for 10–15 min in Slowfade antifade reagent (Molecular Probes). Preparations were mounted onto glass slides in Slowfade antifade under a coverslip. Images of the same NMJs studied for Ca²⁺ imaging were collected under immersion oil with a 40× lens (Nikon), using the Bio-Rad 600 confocal microscope.

Statistics. Results are presented as mean ± SEM. A Student paired t test was used when a treatment was performed on the same cell (control vs drug), and an ANOVA was used for comparison between different groups. Both tests were used at a confidence level of 95% (α = 0.05).

RESULTS

Identification of PSCs at the mammalian NMJ

The validity of the study of synapse–glia interactions at the mammalian NMJ relies on our ability to identify PSCs. However, unlike PSCs at the frog NMJ, mammalian PSCs are not so easily recognized in bright-field illumination. Mammalian NMJs have an oval shape and are more compact than amphibian NMJs. They terminate in a slight elevation of the muscle fiber, just after the myelin sheath where the nerve terminal forms a compact junction (Salpeter, 1987). The identification of PSCs was achieved based on these characteristics.

We first confirmed that these criteria of identification were reliable by labeling PSCs using an antibody against the calcium-binding protein S-100, which is used as a general marker for Schwann cells (Son and Thompson, 1995a). First, cells were identified by transmitted light according to the criteria indicated above, and preparations were loaded with the permeant fluorescent Ca²⁺ indicator fluo-3 AM (Fig. 1A). A brief local application of muscarine increased Ca²⁺ fluorescence in the cell body of the presumed PSCs (Fig. 1B) that subsequently decreased back to baseline (see also Figs. 3 and 4). The nerve–muscle preparation was then processed for immunohistochemistry, and the presence of S-100 was revealed. The same NMJ was then found, and an image of S-100 labeling was taken using the confocal microscope. As shown in Figure 1C, the same cells identified as PSCs corresponding to our criteria and that showed a Ca²⁺ response induced by muscarine were also labeled by S-100. These results indicate that they were Schwann cells at the NMJ, and hence, that the criteria used to identify PSCs were appropriate. Also, the myelinating Schwann cells that were also loaded with fluo-3 were also labeled by S-100 (data not shown).

Nerve-evoked Ca²⁺ responses in perisynaptic Schwann cells

The sensitivity of mouse PSCs to synaptic transmission and transmitter release was established by monitoring changes in intracellular Ca²⁺ during synaptic activity induced by repetitive stimulation of the motor nerve. As shown in Figure 2A, a train of stimuli (50 Hz, 30 sec) triggered an increase in fluorescence. This was observed in all PSCs tested (n = 10) with a mean of 157 ± 15% ΔF/F. This elevation in intracellular Ca²⁺ occurred, on average, with a delay to onset of 1.5 sec after the beginning of the stimulation, and the time to reach the maximum amplitude was 15 ± 2 sec. Ca²⁺ responses decayed to baseline in 82.2 ± 12 sec, which occurred well after the interruption of nerve stimulation. In addition, in all PSCs studied, the initial Ca²⁺ response was always followed by a second smaller elevation that occurred after a partial recovery of the initial response. Nerve stimulation at lower frequencies (10, 15, or 25 Hz) resulted in smaller Ca²⁺ responses than the ones obtained with stimulation at 50 Hz (data not shown). As seen in Figure 2A, a second train of stimuli given 20 min after the first one (break in the x-axis) hardly elicited a
Ca²⁺ response. The mean amplitude of the response was only 14 ± 15% ΔF/F (n = 5). These results indicate that PSCs detect and are sensitive to synaptic activity. In contrast to the PSCs, and although loaded with fluo-3, no Ca²⁺ responses were detected from the myelinating Schwann cells present at the last myelinated segment of the motor nerve (data not shown).

Figure 1. Identification of PSCs at the mammalian NMJ. A, False color confocal image of a mammalian NMJ loaded with the fluorescent Ca²⁺ indicator fluo-3 AM. The range of false colors reflects the level of Ca²⁺ level, where blue indicates a low level of Ca²⁺, and green-red indicates a high level. Two PSCs (arrows) at rest before local application of muscarine. B, The same PSCs as in A, after application of muscarine (20 μM). Note the rise in fluorescence induced by muscarine. C, Labeling of PSCs of the same NMJ with anti-S100, a calcium binding protein used as a general marker for Schwann cells. Note that the two cells that responded to muscarine were also labeled by the S100 antibody. Scale bar, 20 μm.

Figure 2. Nerve-evoked Ca²⁺ responses in PSCs. Ca²⁺ responses obtained using the fluorescent Ca²⁺ indicator fluo-3 AM and monitored using a Bio-Rad 600 confocal microscope. Blue indicates low level of calcium, and red indicates a high level. A, Changes in fluorescence (%ΔF/F) before, during (bar), and after motor nerve stimulation (50 Hz, 30 sec). Note that relative increase in Ca²⁺ fluorescence to a second train of stimuli applied 20 min later (break in the x-axis) was much reduced. B, Ca²⁺ response in the same PSC as in A, before (1), during (2), and after (3–5) transmitter release induced by repetitive stimulation of the motor nerve (50 Hz, 30 sec). The corresponding time at which each image was taken is illustrated on the graph in A with the corresponding number of the figure above. C, Changes in fluorescence (%ΔF/F) before, during (bar), and after motor nerve stimulation (50 Hz, 30 sec) in the presence of α-conotoxin MVIIC (1 μM). Note that the increase in the intensity of the fluorescence was strongly reduced when transmitter release was blocked by α-conotoxin MVIIC, a P/Q-type Ca²⁺ channel blocker. Different preparation than A and B. D, Mean ± SEM of amplitude of Ca²⁺ responses in PSCs elicited by nerve stimulation in the absence and in the presence of α-conotoxin MVIIC (p = 0.001; Student’s paired t test). Scale bar: B, 10 μm.
Is transmitter release required to elicit Ca$^{2+}$ responses?

The response of PSCs to synaptic activity could be attributable to an increase of extracellular K$^+$ ions during high-frequency stimulation that would depolarize the cell and open voltage-gated Ca$^{2+}$ channels (MacVicar, 1984; Newman, 1986; Barres et al., 1990; Robitaille et al., 1996) or to the release of neurotransmitters (Jahromi et al., 1992; Reist and Smith, 1992; Robitaille, 1995). To distinguish between the two possibilities, the Ca$^{2+}$-dependent release of neurotransmitters was prevented by blocking the Ca$^{2+}$ channels that trigger this process. If transmitter release is required to elicit Ca$^{2+}$ responses observed in PSCs, these should be reduced in the presence of the Ca$^{2+}$ channel blocker. However, Ca$^{2+}$ responses in PSCs should remain unchanged if they are caused by K$^+$ accumulation. Transmitter release was blocked using the toxin ω-conotoxin MVIIIC (ω-CgTX), which binds irreversibly to P/Q-type Ca$^{2+}$ channel. It was used at concentrations known to completely block evoked transmitter release at the mouse NMJ (Katz et al., 1996). After blockade of transmitter release by ω-CgTX (1 μM; data not shown), the nerve-evoked Ca$^{2+}$ signal in PSCs was on average 9 ± 2% ΔF/ΔF₀ (n = 6) (Fig. 2C). This represents a reduction of 94% in the size of the Ca$^{2+}$ responses in comparison to the ones obtained in the absence of ω-CgTX MVIIIC (Fig. 2D). Thus, substances released by the nerve terminal during evoked activity are necessary for the induction of Ca$^{2+}$ responses in PSCs, suggesting that PSCs are sensitive to transmitters released during synaptic activity. The residual Ca$^{2+}$ response in PSCs in the presence of ω-CgTX MVIIIC is likely attributable to the depolarization of the cell caused by K$^+$ accumulation during synaptic activity (Jahromi et al., 1992; Robitaille et al., 1996).

Can ACh trigger Ca$^{2+}$ responses at the mammalian NMJ?

Our data indicate that neurotransmitter substances released during synaptic activity are necessary to elicit fully developed Ca$^{2+}$ responses in PSCs. As primary candidates, ACh and ATP appear particularly important because ACh is the main neurotransmitter at this synapse and ATP is coreleased at an equimolar concentration (Silinsky, 1975; Smith, 1991). If substances such as the cotransmitters ATP and ACh released by the active nerve terminal induce PSC Ca$^{2+}$ signals, then application of these substances alone should mimic the effects induced by stimulation of the motor nerve.

The ability of PSCs to respond to ACh applied locally was examined first. ACh was applied directly by micropressure (5–10 PSI) to selected PSCs. As shown in Figure 3A, local application of ACh (20 μM) induced a Ca$^{2+}$ response in all PSCs tested (350 ± 18% ΔF/ΔF₀; n = 10). Because the Ca$^{2+}$ response obtained by stimulation of the motor nerve decreased during consecutive trains of stimuli (Fig. 2A), we wondered whether the Ca$^{2+}$ responses induced by local applications of ACh would also be susceptible to rundown. Indeed, a second application of ACh evoked responses that were significantly smaller than the first ones (249 ± 19% ΔF/ΔF₀ vs 357 ± 30% ΔF/ΔF₀ for control; n = 5; p = 0.0006; Student’s paired t test) (Fig. 3A, break in the x-axis).

We next investigated whether the rise in intracellular Ca$^{2+}$ triggered by ACh was attributable to an entry of extracellular Ca$^{2+}$ or caused by the release from intracellular stores. To distinguish between the two possibilities, extracellular Ca$^{2+}$ was removed and replaced by a 5 mM Mg$^{2+}$ physiological solution, and the ability of PSCs to respond to locally applied ACh was tested again. As shown in Figure 3B, on all PSCs tested, Ca$^{2+}$ responses were elicited by a local application of ACh in 0 Ca$^{2+}$/5 Mg$^{2+}$. 20 min after a first application in normal Ca$^{2+}$, indicating that ACh induced the release of Ca$^{2+}$ from internal stores. Surprisingly, the size of Ca$^{2+}$ responses induced by ACh in 0 Ca$^{2+}$/5 Mg$^{2+}$ was significantly larger than the size of the responses obtained in the presence of extracellular Ca$^{2+}$ (421 ± 15% ΔF/ΔF₀ vs 343 ± 22% ΔF/ΔF₀ for control; n = 5; p = 0.001; Student’s paired t test) (Fig. 3B). This indicates that, although the contribution of internal stores is essential for the induction of Ca$^{2+}$ responses by ACh, the presence of extracellular Ca$^{2+}$ downregulates the activation of the release of Ca$^{2+}$ from internal stores by ACh (Fig. 3C).
Muscarinic receptors mediate ACh effects

For a better understanding of the role of glial cells at the synapse, it is necessary to further characterize the cellular receptors and functions of these glial cells. To determine the type of cholinergic receptors involved, we first used α-BuTx, an antagonist of the nicotinic ACh receptors (AChRs). If Ca\(^{2+}\) responses to ACh were caused by nicotinic AChRs, they should be blocked by the nicotinic cholinergic antagonist α-BuTx. To test the effectiveness of α-BuTx (20 μM), preparations were perfused with the antagonist for at least 30 min before local application of ACh on PSCs was performed. This concentration blocks completely the activity of the nicotinic postsynaptic receptors at the frog NMJ (Jahromi et al., 1992). As shown in Figure 4, A and B, the ability of PSCs to respond to ACh was not affected by the presence of α-BuTx. Indeed, Ca\(^{2+}\) responses to ACh were still obtained in the presence of the toxin in all cells tested, and the size of the Ca\(^{2+}\) responses (317 ± 13% ΔF/F; n = 5) was not significantly different from the control responses obtained on the same cells before the application of α-BuTx (Student’s paired t test; p > 0.05).

The lack of effect of the nicotinic antagonist suggests that PSC AChRs are not nicotinic but rather, that the Ca\(^{2+}\) response induced by ACh was likely because of the activation of muscarinic receptors. If this were the case, muscarinic antagonists such as atropine should prevent Ca\(^{2+}\) responses to ACh. As shown in Figure 4B, the presence of atropine almost completely abolished Ca\(^{2+}\) responses elicited by ACh in all cells tested (n = 15). The mean size of the responses was only 14 ± 6% ΔF/F, which is significantly smaller than the responses obtained from the same cells by a first application of ACh (Student’s paired t test; p = 0.001).

We next investigated the ability of the muscarinic agonist muscarine to evoke Ca\(^{2+}\) responses. Local application of muscarine (20 μM) induced a Ca\(^{2+}\) response in all PSCs studied. A second application 20 min later (break in the x-axis) induced smaller changes in fluorescence. D, Changes in fluorescence caused by the local application of muscarine (20 μM) in the presence (solid line) and in the absence (dashed line) of external Ca\(^{2+}\) for the same PSCs. Ca\(^{2+}\) was replaced by 5 mM Mg\(^{2+}\). Note that muscarine induced larger changes in fluorescence in the absence of external Ca\(^{2+}\). E, Comparison of mean ± SEM of Ca\(^{2+}\) responses induced by a first and a second application of muscarine (left) and application of muscarine in presence and absence of external Ca\(^{2+}\) (right).
Can ATP and adenosine trigger Ca\(^{2+}\) responses at the mammalian NMJ?

At the frog NMJ, application of ATP to PSCs induces the release of Ca\(^{2+}\) from internal stores (Jahromi et al., 1992; Robitaille, 1995) because of the activation of P\(_{2\alpha}\) and P\(_{2\gamma}\) receptors (Robitaille, 1995). To examine the involvement of endogenous purines in the modulation of mammalian PSCs in situ and to test whether ATP and adenosine can activate PSCs, changes in intracellular Ca\(^{2+}\), induced by local application of ATP and adenosine were monitored.

ATP (20 \(\mu\)M) induced Ca\(^{2+}\) responses in all PSCs tested (299 \(\pm\) 30% \(\Delta F/F\); \(n = 41\)) (Fig. 5A), whereas local applications of adenosine (20 \(\mu\)M) resulted in a rise of fluorescence of 441 \(\pm\) 51% \(\Delta F/F\) also in all PSCs tested (\(n = 19\)) (Fig. 5B). A second application of ATP evoked responses that were significantly smaller than the first response (207 \(\pm\) 30% \(\Delta F/F\) vs 246 \(\pm\) 38% \(\Delta F/F\) for control; \(n = 8\); \(p = 0.0004\)) (Fig. 5A). Similarly, a second application of adenosine evoked responses that were significantly smaller than the first response (314 \(\pm\) 52% \(\Delta F/F\) vs 450 \(\pm\) 77% \(\Delta F/F\); \(n = 11\); \(p = 0.0004\); Student’s paired \(t\) test) (Fig. 5B).

In most glial cells (Salter and Hicks, 1994), including PSCs at the amphibian NMJ (Jahromi et al., 1992; Robitaille, 1995), purinergic agonists elicit an elevation in intracellular Ca\(^{2+}\) that involves the release of Ca\(^{2+}\) from internal stores. This possibility was tested by replacing extracellular Ca\(^{2+}\) ions by Mg\(^{2+}\) (5 mM). As shown in Figure 5C, Ca\(^{2+}\) responses induced by local application of ATP still induced Ca\(^{2+}\) responses in all mammalian PSCs tested, even in the absence of Ca\(^{2+}\) in the solution, pointing toward a critical role of Ca\(^{2+}\) internal stores in the production of the Ca\(^{2+}\) responses. On average, the amplitude of the Ca\(^{2+}\) responses was 461 \(\pm\) 98% \(\Delta F/F\). However, similar to ACh-induced Ca\(^{2+}\) responses, local application of ATP in absence of extracellular Ca\(^{2+}\) evoked Ca\(^{2+}\) responses that were significantly larger than the ones evoked by a first application of ATP on the same cells (273 \(\pm\) 42% \(\Delta F/F\) for control; \(n = 11\); \(p = 0.02\); Student’s paired \(t\) test) (Fig. 5C). Similarly, adenosine also caused Ca\(^{2+}\) responses in absence of external Ca\(^{2+}\) that were significantly larger than those obtained in the presence of Ca\(^{2+}\) (497 \(\pm\) 113% \(\Delta F/F\); \(n = 11\); \(p = 0.0004\); Student’s paired \(t\) test) (Fig. 5D).
72% ΔF/F vs 428 ± 6% ΔF/F; n = 8; p = 0.001; Student’s paired t test) (Fig. 5D). These results indicate that PSCs at the mammalian NMJ detect adenosine and ATP and that these substances induce the release of Ca\(^{2+}\) from internal stores via a mechanism that is negatively regulated by external Ca\(^{2+}\) (Fig. 5E,F).

**Are purinergic receptors present on PSCs?**

To verify the presence of adenosine and ATP receptors on the mammalian PSCs, the ability of these agonists to evoke Ca\(^{2+}\) responses in PSCs was tested in the presence of antagonists for P2 and P1 receptors. Suramin, a nonselective P2 receptor antagonist was first tested. Ca\(^{2+}\) responses induced by ATP should be abolished in the presence of this antagonist if they are mediated by the activation of P2 receptors. After eliciting a Ca\(^{2+}\) response by ATP on PSCs, the preparations were perfused with suramin (100 μM), and the ATP was applied locally on the same cells. As shown in Figure 6, Ca\(^{2+}\) responses were still induced by local applications of ATP (20 μM) (289 ± 34% ΔF/F; n = 6) (Fig. 6A), and the average amplitude of the responses was not significantly different from responses evoked by the same cells before application of the antagonist (Student’s paired t test; p = 0.239). It is unlikely that the lack of effect was attributable to the inability of suramin to block P2 receptors because the same solution blocked ATP-induced responses at PSCs of amphibian NMJs (Robitaille, 1995). Hence, this result suggests that ATP raised intracellular Ca\(^{2+}\) by mechanisms that do not activate P2 receptors.

However, Ca\(^{2+}\) responses evoked by adenosine were almost completely abolished in the presence of CPT (10 μM), an A1 receptor antagonist (32 ± 6% ΔF/F; n = 5) (Fig. 6B). This value was significantly smaller than the responses obtained on the same cells before the application of the antagonist in the perfusate (Student’s paired t test; p = 0.001). This suggests that the adenosine-induced Ca\(^{2+}\) responses were mediated by the activation of A1 receptors.

Ectoenzymes present in the synaptic cleft (Salter et al., 1993; Ziganshin et al., 1993) will cause the dephosphorylation of ATP and lead to adenosine formation. Because adenosine appears as an active substance at the mammalian NMJ and because ATP can be exogenously degraded into adenosine, it is possible that the Ca\(^{2+}\) responses evoked by local applications of ATP were attributable to the indirect activation of A1 receptors. This possibility was tested by locally applying ATP in the presence of CPT. As shown in Figure 6C, the presence of CPT (10–20 μM) in the perfusion did not block Ca\(^{2+}\) responses induced by ATP. The average amplitude of the Ca\(^{2+}\) responses was 384 ± 86% ΔF/F (n = 7), which is not significantly different from control (ANOVA; p > 0.05). These results suggest that the action of ATP was not mediated by an indirect activation of A1 receptors and, hence, would be produced by other ATP-dependent mechanisms.

**Endogenous ACh and adenosine mediate nerve-evoked Ca\(^{2+}\) responses in PSCs**

Our results indicate that functional muscarinic and A1 adenosine receptors are present on PSCs of the mammalian NMJ. We next wondered whether ACh and adenosine released by the presynaptic terminal during synaptic activity could mediate the activation of PSCs and the triggering of the nerve stimulation-evoked Ca\(^{2+}\) responses in PSCs. Because nerve-evoked Ca\(^{2+}\) responses decrease during consecutive trains of stimuli (Fig. 2A), the effects of the two different blockers on nerve-evoked Ca\(^{2+}\) responses were tested on different preparations, and each preparation was stimulated only once.

![Figure 6](image-url)

Figure 6. Ca\(^{2+}\) responses of PSCs to local application of ATP and adenosine in presence of purinergic antagonists. A, Changes in fluorescence caused by the local application of ATP before and after (break in the x-axis) a 20 min perfusion with suramin (100 μM), a nonspecific P2 antagonist. Note that the increase in the intensity of fluorescence induced by the local application of ATP was unaffected by the presence of suramin. B, Changes in fluorescence caused by the local application of adenosine before and after a 20 min perfusion with CPT (10 μM), a nonspecific P2 antagonist. Note that the increase in the intensity of fluorescence induced by the local application of ATP was unaffected by the presence of suramin. C, Changes in fluorescence in four PSCs of an NMJ caused by the local application of ATP in the presence of the A1 receptor antagonist CPT (20 μM). Inset, False color confocal images of the four PSCs at rest (Rest), at the peak of the Ca\(^{2+}\) response (Peak), and after recovery (Recov). Note that the increase in the intensity of fluorescence induced by the local application of ATP was unaffected by the presence of CPT. Scale bar, 10 μm.

We first examined the impact of blockade of ACh receptors on nerve-evoked Ca\(^{2+}\) responses. The blockade of ACh receptors with atropine (20 μM) greatly reduced the amplitude of Ca\(^{2+}\) responses induced by stimulation of the motor nerve to 33 ± 3% ΔF/F (n = 5) (Fig. 7A). This value is significantly smaller than the
We present here direct evidence that mammalian PSCs in situ respond to the release of neurotransmitters induced by stimulation of the motor nerve, as indicated by an elevation of intracellular Ca\(^{2+}\) level. This elevation originated mainly from the release of Ca\(^{2+}\) from internal stores and muscarinic and adenosine A1 receptor antagonists together. As shown in Figure 7C, Ca\(^{2+}\) responses were primarily reduced in the presence of the two antagonists in which the mean Ca\(^{2+}\) response was only 22 ± 8% (N = 4; n = 8). This value is significantly different from the mean response obtained in control condition (i.e., without any antagonist; one-way ANOVA, p < 0.05). The remaining Ca\(^{2+}\) elevation is consistent with the Ca\(^{2+}\) response observed when transmitter release was blocked using ω-CgTx MVIIIC, possibly because of the depolarization of PSCs by K\(^+\) accumulation during repetitive nerve stimulation. The inability to elicit Ca\(^{2+}\) responses was not caused by a lack of cell responsivity because local application of ATP (brake in x-axis) induced Ca\(^{2+}\) responses in all cells tested (N = 3; n = 7).

**DISCUSSION**

We next tested the impact of the A1 antagonist CPT on Ca\(^{2+}\) responses evoked by stimulation of the motor nerve. As shown in Figure 7B, Ca\(^{2+}\) responses were reduced to 98 ± 6% \(\Delta F/F\) (n = 6) (Fig. 7B) in the presence of CPT. Although the reduction in the size of Ca\(^{2+}\) responses was more modest than the effect of the muscarinic antagonist, the size of the Ca\(^{2+}\) responses was significantly smaller than the control ones (one-way ANOVA; p < 0.05) (Fig. 7B, gray zone). This indicates that the activation of adenosine receptors is important to generate the Ca\(^{2+}\) responses induced by nerve-evoked transmitter release. The partial blockade of the responses was not attributable to the inefficacy of CPT to antagonize the A1 receptors because local application of adenosine on the same cells did not induce any Ca\(^{2+}\) responses (data not shown).

Finally, we tested the impact on Ca\(^{2+}\) responses of both the muscarinic and A1 receptor antagonists together. As shown in Figure 7C, Ca\(^{2+}\) responses were primarily reduced in the presence of the two antagonists in which the mean Ca\(^{2+}\) response was only 22 ± 8% (N = 4; n = 8). This value is significantly different from the mean response obtained in control condition (i.e., without any antagonist; one-way ANOVA, p < 0.05). The remaining Ca\(^{2+}\) elevation is consistent with the Ca\(^{2+}\) response observed when transmitter release was blocked using ω-CgTx MVIIIC, possibly because of the depolarization of PSCs by K\(^+\) accumulation during repetitive nerve stimulation. The inability to elicit Ca\(^{2+}\) responses was not caused by a lack of cell responsivity because local application of ATP (brake in x-axis) induced Ca\(^{2+}\) responses in all cells tested (N = 3; n = 7).

**Cholinergic receptor on the mammalian PSCs**

When expressed by glial cells, including PSCs of the amphibian NMJ (Robitaille et al., 1997), cholinergic receptors have always been determined to be of the muscarinic type, mainly linked to internal Ca\(^{2+}\) stores regulated by an inositol 1,4,5-trisphosphate receptor (Hamprecht, 1986). Here we show that PSCs of the mammalian NMJ are no exception and that the cholinergic receptors are of the muscarinic type as indicated by the lack of effects of nicotine antagonists and by the ability of muscarine to induce Ca\(^{2+}\) responses, whereas atropine blocked ACh-induced responses. The blockade of Ca\(^{2+}\) responses by the nonspecific muscarinic antagonist atropine also indicates that PSCs of the mammalian NMJ, unlike the ones at the amphibian NMJ, express a typical muscarinic pharmacology (Robitaille et al., 1997). Moreover, muscarinic-dependent Ca\(^{2+}\) responses in glial cells also
share another characteristic, that is, the rundown of the response after repetitive applications of cholinergic agonists (Dave et al., 1991; Jahromi et al., 1992; Robitaille et al., 1997).

### Purinergic receptors on the mammalian PSCs

It is now established that ATP is released during synaptic transmission (Smith, 1991; Zimmermann, 1994) and is involved in intercellular signaling in various systems (Sneddon and Burnstock, 1984; Benham, 1989; Salter and Hicks, 1994; Zimmermann, 1994; Lyons et al., 1995). Adenosine and ATP receptors are widely distributed (Burnstock, 1990; Salter et al., 1993) and are present on several types of glial cells such as cultured astrocytes (Lai and Wong, 1991; Salter and Hicks, 1994), glioma cells (Chueh et al., 1994), microglial cells (Walz et al., 1993), and glial cells of the rat optic nerve (Kriegler and Chiu, 1993). Moreover, ATP appears necessary for the spread of Ca\(^{2+}\) waves in a synaptium of astrocytes (Guthrie et al., 1999).

The results of the present study suggest that PSCs possess adenosine receptors of A1 type, as indicated by the blockade of adenosine-induced Ca\(^{2+}\) responses by CPT, an A1 receptor antagonist. However, and somewhat unexpectedly, Ca\(^{2+}\) responses induced by local applications of ATP were not blocked by suramin, a nonspecific P2 receptor antagonist indicating that the effects observed with ATP were not attributable to the activation of P2 receptors. In addition, adenosine released during synaptic activity triggers Ca\(^{2+}\) responses, as indicated by the significant reduction of Ca\(^{2+}\) responses elicited by nerve stimulation in the presence of CPT. However, at the amphibian NMJ, ATP released during synaptic transmission activates PSCs, whereas endogenous adenosine had no apparent effect on nerve-evoked Ca\(^{2+}\) responses (Robitaille, 1995).

Because of the presence of ecto-ATPases (Zimmermann, 1994), one possibility might have been that ATP-induced responses were caused by the indirect activation of adenosine receptors as a consequence of ATP degradation into adenosine. However, our results are not consistent with this possibility because ATP-induced Ca\(^{2+}\) responses persisted even in the presence of the adenosine receptor antagonist. An alternative possibility might be that ATP acted via ecto-protein-kinases because they are known to be located on the surface of several cells (Zimmermann, 1994) and that they require the presence of Mg\(^{2+}\) ions (Ehrlich et al., 1986), conditions in which our experiments were performed.

### Glial cells detect neuronal activity and synaptic transmission

Glial cells at CNS synapses detect the release of neurotransmitters induced by neuronal activity (Porter and McCarthy, 1996, 1997; Carmignoto et al., 1998; Grosche et al., 1999). This does not require the activation of a complex multicellular network because similar results were obtained at the frog NMJ where PSCs detect and are modulated by synaptic activity generated by a single synapse (Jahromi et al., 1992; Georgiou et al., 1994, 1999; Robitaille, 1995; Robitaille et al., 1997; Bourque and Robitaille, 1998).

The data presented here demonstrate that mammalian PSCs can also detect synaptic activity and that a number of Ca\(^{2+}\)-dependent cascades of events are likely triggered in PSCs as a consequence of the Ca\(^{2+}\) elevations. This observation is consistent with the properties reported for all other synapses at which synapse–glia interactions have been studied (Araque et al., 1999; Castonguay et al., 2001). Indeed, synaptic activity will trigger an elevation of intracellular Ca\(^{2+}\) in PSCs as a result of its release from internal stores. Moreover, this activation is frequency dependent and is regulated by the main neurotransmitters present at the synapse, regulating G-protein-coupled receptors. Hence, the results obtained at the mouse NMJ further support the concept that synapse–glia interactions are ubiquitous at chemical synapses where glial processes are present.

Because synapse–glia interactions are dependent on the release of neurotransmitters by chemical synapses (Araque et al., 1999; Castonguay et al., 2001), it is expected that the activity of perisynaptic glial cells will be governed by the main neurotransmitters found at the synapses with which they are associated. In the present study, it is shown that PSCs activation requires the release of neurotransmitter, as shown by the blockade of transmitter release with α-CgTx MVIIIC. In particular, ACh released during synaptic transmission modulates the activity of the PSCs because Ca\(^{2+}\) responses were almost completely abolished in the presence of the muscarinic receptor antagonist atropine. Consistent with this observation and similar to ACh-induced Ca\(^{2+}\) responses, we found that Ca\(^{2+}\) responses induced by evoked transmitter release also showed a pronounced rundown when elicited by repetitive trains of stimuli. This property was also present at PSCs of the amphibian NMJ (Jahromi et al., 1992) where a large part of this rundown was shown to be caused by endogenously released substance P that activates NK-1 receptors leading to reduced efficacy of purinergic and muscarinic receptor systems (Bourque and Robitaille, 1998).

### Functional differences of mammalian and amphibian PSCs

Although mammalian and amphibian PSCs share a number of properties, there are a few major functional differences (Table 1).

**Table 1. Comparison of the properties of PSCs at mammalian and amphibian NMJs**

<table>
<thead>
<tr>
<th></th>
<th>Cholinergic receptors</th>
<th>Adenosine receptors</th>
<th>ATP receptors</th>
<th>Ca(^{2+}) mechanisms</th>
<th>Ca(^{2+}) regulation</th>
<th>Oscillation</th>
<th>Rundown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouse NMJ</td>
<td>Muscarinic</td>
<td>A(_1)</td>
<td>No</td>
<td>Internal stores</td>
<td>Negative regulation</td>
<td>Yes</td>
<td>Very steep</td>
</tr>
<tr>
<td>Frog NMJ</td>
<td>Muscarinic, not M1–M5</td>
<td>A(_1)</td>
<td>P(_{2_X})</td>
<td>Internal stores; IP(_{3R})</td>
<td>No</td>
<td>No</td>
<td>Gradual</td>
</tr>
</tbody>
</table>

Comparison of the functional similarities and differences of mouse and frog PSCs concerning the type of receptors, Ca\(^{2+}\) regulatory mechanisms, and responses to synaptic transmission evoked by nerve stimulation.
or stimulation of the motor nerve. This was not observed at amphibian PSCs where muscarine induced a single, large increase in Ca\(^{2+}\) (Jahromi et al., 1992; Robitaille, 1995; Robitaille et al., 1997). Moreover, the Ca\(^{2+}\) responses appear negatively regulated by external Ca\(^{2+}\) at mammalian but not at amphibian PSCs. The presence of the negative regulation by external Ca\(^{2+}\) is consistent with a calcium-dependent/calcium-release phenomenon that would only be present in mammalian PSCs. This negative Ca\(^{2+}\) regulation may contribute to the genesis of the repeated Ca\(^{2+}\) elevations observed in PSCs and to the more pronounced rundown of consecutive Ca\(^{2+}\) responses elicited at mammalian PSCs. These differences will have major impacts on the functions of PSCs because different types of receptors will have different regulatory actions resulting in the production of different second messengers. Also, the different Ca\(^{2+}\) regulation in PSCs will result in the induction of different Ca\(^{2+}\)-dependent mechanisms. Hence, although amphibian and mammalian NMJs show fundamental synapse–glia interactions, the properties of these interactions differ.

**Properties of synapse–glia interactions as a function of synapse identity**

Based on our data, we propose that, although synapse–glia interaction is a common phenomenon at chemical synapses associated with glial cells, the nature and the extent of the interactions will be fine-tuned according to the properties and function of the synapse with which they are associated. Because perisynaptic glial cells are involved in the modulation of synaptic transmission (Parpura et al., 1994; Robitaille, 1998; Araque et al., 1999; Castonguay et al., 2001), their involvement in the regulation of synaptic functions will be tuned according to their functional properties and, hence, adjusted to their synaptic environment. This adaptation to the synaptic environment is essential for perisynaptic glial cells to actively and effectively modulate synaptic efficacy and neuronal activity. Thus, it will be critical to establish the properties of the regulation of synaptic efficacy by PSCs at the mouse NMJ and determine how the different properties of PSCs at this synapse will influence their regulation of synaptic activity. Moreover, because of the importance of PSCs in the regulation of neuronal activity and synaptic functions will be tuned according to their functional properties and, hence, adjusted to their synaptic environment. Thus, it will be critical to establish the properties of the regulation of synaptic efficacy by PSCs at the mouse NMJ and determine how the different properties of PSCs at this synapse will influence their regulation of synaptic activity. Moreover, because of the importance of PSCs in the regulation of neuronal activity and synaptic functions will be tuned according to their functional properties and, hence, adjusted to their synaptic environment.

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